## **AMENDMENTS TO THE CLAIMS**

## Claims 1-4 (cancelled)

- 5. (currently amended) A method of determining the identification of a nucleotide at a detection position in a target sequence comprising a first target domain adjacent to said detection position and a second target domain comprising said detection position, wherein said method comprises:
- a) hybridizing a first ligation probe to said first target domain, said first ligation probe comprising:
  - i) a first portion comprising an upstream universal priming site (UUP); and
  - ii) a second portion comprising a first target-specific sequence comprising a first base at an interrogation position; and
- b) hybridizing a second ligation probe to said second target domain, said second ligation probe comprising:
  - i) a third portion comprising a downstream universal priming site (DUP); and
  - ii) a fourth portion comprising a second target-specific sequence;

wherein if said first base is perfectly complementary to said nucleotide at said detection position, a ligation complex is formed, and

wherein at least one of said first and second ligation probes comprises a fifth portion comprising an adapter sequence;

- c) removing non-hybridized probes;
- d) providing a ligase that ligates said first and second ligation probes to form a ligated probe;
- e) amplifying said ligated probe using said UUP and said DUP to generate a plurality of amplicons;
  - f) contacting said amplicons with an array of capture probes; and
  - g) determining the nucleotide at said detection position.

## Claims 6-8 (cancelled)

- 9. (previously amended) A method according to claim 5, 26, 32 and 33 wherein said removing comprises:
- a) enzymatically adding a binding ligand to said target sequence to form a target sequence comprising said binding ligand;

- b) binding a hybridization complex comprising said target sequence comprising said binding ligand to a binding partner immobilized on a solid support;
  - c) washing away unhybridized probes; and
  - d) eluting said probes off said solid support.
- 10. (previously amended) A method according to claim 5, 26, 32, or 33 wherein said removing is done using a double-stranded specific moiety.
- 11. (original) A method according to claim 10 wherein said double-stranded specific moiety is an intercalator attached to a support.
- 12. (previously amended) A method according to claim 11 wherein said support is a bead.
- 13. (**previously amended**) A method according to claim 5, 26, 32, or 33 wherein said amplifying is done by:
  - a) hybridizing a first universal primer to said UUP;
  - b) providing a polymerase and dNTPs such that said first universal primer is extended;
  - c) hybridizing a second universal primer to said DUP;
- d) providing a polymerase and dNTPs such that said second universal primer is extended; and
  - e) repeating steps a) through d).
- 14. (previously amended) A method according to claim 5, 26, 32, or 33 wherein said array comprises:
  - a) a substrate with a patterned surface comprising discrete sites; and
- b) a population of microspheres comprising at least a first subpopulation comprising a first capture probe and a second subpopulation comprising a second capture probe.
- 15. (original) A method according to claim 14 wherein said discrete sites comprise wells.
- 16. (**previously amended**) A method according to claim 14 wherein said substrate comprises a fiber optic bundle.

Claims 17-18 (canceled)

- 19. (previously amended) A method according to claim 5 or 32, further comprising providing a support on which the target sequence is immobilized.
- 20. (original) A method according to claim 19, wherein said non-hybridized probes are removed without removing said target sequence from said support.

- 21. (previously amended) A method according to claim 5 or 32, further comprising attaching said target sequence to a support.
- 22. (currently amended) A method according to claim 21, wherein said target sequence is attached to said support by a method selected from the group consisting of labeling said target sequence with a functional attachment moiety capable of binding to that binds said support and interacting said functional attachment moiety with said support, absorption of said target sequence on a charged support said support wherein said support comprises charged groups, direct chemical attachment of said target sequence to said support and photocrosslinking said target sequence to said support.
- 23. (previously amended) A method according to claim 9, wherein said support is selected from the group consisting of paper, plastic and tubes.

Claims 24-25 (cancelled)

- 26. (currently amended) A method of determining the identification of a nucleotide at a detection position in a target sequence comprising a first target domain adjacent to said detection position and a second target domain comprising said detection position, wherein said method comprises:
  - a) providing a support on which the target sequence is immobilized;
- b) hybridizing a first ligation probe to said first target domain, said first ligation probe comprising:
  - i) a first portion comprising an upstream universal priming site (UUP); and
  - ii) a second portion comprising a first target-specific sequence comprising a first base at an interrogation position; and
- c) hybridizing a second ligation probe to said second target domain, said second ligation probe comprising:
  - i) a third portion comprising a downstream universal priming site (DUP); and
- ii) a fourth portion comprising a second target-specific sequence; wherein if said first base is perfectly complementary to said nucleotide at said detection position, a ligation complex is formed, and wherein at least one of said first and second ligation probes comprises a fifth portion comprising an adapter sequence;
  - d) removing non-hybridized probes;

- e) providing a ligase that ligates said first and second ligation probes to form a ligated probe;
- f) amplifying said ligated probe using said UUP and said DUP to generate a plurality of amplicons;
  - g) contacting said amplicons with an array of capture probes; and
  - h) determining the nucleotide at said detection position.

## Claims 27-29 (cancelled)

- 30. (previously added) A method according to claim 9 wherein said solid support is a bead.
- 31. (previously added) A method according to claim 26 wherein said non-hybridized probes are removed without removing said target sequence from said support.
- 32. (currently amended) A method of determining the identification of a nucleotide at a detection position in a target sequence comprising a first target domain comprising said detection position and a second target domain adjacent to said detection position, wherein said method comprises:
- a) hybridizing a first ligation probe to said first target domain, said first ligation probe comprising:
  - i) a first portion comprising an upstream universal priming site (UUP);
  - ii) a second portion comprising a first target-specific sequence; and
  - iii) an interrogation position that is complementary to said detection position; and
- b) hybridizing a second ligation probe to said second target domain, said second ligation probe comprising:
  - i) a third portion comprising a downstream universal priming site (DUP); and
- ii) <u>a fourth portion comprising</u> a second target-specific sequence; whereby if said interrogation position of said first probe is perfectly complementary to said nucleotide at said detection position, a ligation complex is formed, <u>and</u> wherein at least one of said first and second ligation probes comprises <u>a fifth portion comprising</u> an adapter sequence;
  - c) removing non-hybridized probes;
- d) providing a ligase that ligates said first and second ligation probes to form a ligated probe;
- e) amplifying said ligated probe using said UUP and said DUP to generate a plurality of amplicons;

- f) contacting said amplicons with an array of capture probes; and
- g) determining the nucleotide at said detection position.
- 33. (currently amended) A method of determining the identification of a nucleotide at a detection position in a target sequence comprising a first target domain comprising said detection position and a second target domain adjacent to said detection position, wherein said method comprises:
  - a) providing a support on which the target sequence is immobilized;
  - b) hybridizing a first ligation probe to said first target domain, said first ligation probe comprising:
    - i) a first portion comprising an upstream universal priming site (UUP);
    - ii) a second portion comprising a first target-specific sequence; and
    - iii) an interrogation position; and
  - c) hybridizing a second ligation probe to said second target domain, said second ligation probe comprising:
    - i) a third portion comprising a downstream universal priming site (DUP); and
- ii) a fourth portion comprising a second target-specific sequence; whereby if said interrogation position of said first probe is perfectly complementary to said nucleotide at said detection position, a ligation complex is formed, and wherein at least one of said first and second ligation probes comprises a fifth portion comprising an adapter sequence;
  - d) removing non-hybridized probes;
  - e) providing a ligase that ligates said first and second ligation probes to form a ligated probe;
- f) amplifying said ligated probe using said UUP and said DUP to generate a plurality of amplicons;
  - g) contacting said amplicons with an array of capture probes; and
  - h) determining the nucleotide at said detection position.
- 34. (previously added) A method according to claim 15, wherein said substrate comprises a fiber optic bundle.

- 35. (New) The method according to claim 22, wherein said target sequence is attached to said support by labeling said target sequence with a functional attachment moiety capable of binding to said support and interacting said functional attachment moiety with said support.
- 36. (New) The method according to claim 22, wherein said target sequence is attached to said support by absorption of said target sequence on said support wherein said support comprises charged groups.
- 37. (New) The method according to claim 22, wherein said target sequence is attached to said support by direct chemical attachment of said target sequence to said support.
- 38. (New) The method according to claim 22, wherein said target sequence is attached to said support by photocrosslinking said target sequence to said support.